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I Claim:

- 1. A method for amplifying the presence of an actively respiring microorganisms in a sample comprising contacting the contents of said sample to a nutrient medium containing a predetermined amount of a viability substrate, wherein metabolism of said
- 5 viability substrate by the microorganisms produces a viability marker.
 - 2. The method of claim 1 wherein the microorganisms comprise bacteria.
 - The method of claim 1 wherein the viability substrate is triphenyltetrazolium, nitrotetrazolium blue, iodonitrotetrazolium or dimethylthiazolyldiphenyl tetrazolium.
 - 4. The method of claim 1 wherein the nutrient media contains glucose and NADH.
 - The method of claim 1 wherein the viability marker is a water insoluble marker that accumulates in direct proportion to the number of microorganisms of said sample.
 - A method for detecting an actively respiring microorganisms in a sample comprising:

trapping the microorganisms on a solid filtration membrane; treating the microorganisms according to the method of claim 1; digesting the microorganisms;

contacting secondary antibodies prepared against the primary antibodies

contacting primary antibodies prepared against a substituted formazan with the digested microorganisms to capture said primary antibodies;

20 and conjugated with a detectable marker to captured primary antibodies; and detecting the secondary antibodies that are bound to the captured primary antibodies.

- A method for detecting microorganisms whose presence is amplified by the method of claim 1 comprising:
- 25 digesting the microorganisms by incubation with a lysozyme to form a cellular debris, wherein the viability marker is adsorbed on a surface of the cellular debris:

immobilizing primary antibodies specific for the viability marker on a solid support;

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contacting the digested microorganisms with the immobilized primary antibodies thereby capturing the microorganisms, and

detecting the presence of the viability marker.

- 8. The method of claim 7 wherein the step of detecting comprises:
- 5 contacting the captured digested microorganisms with a reporter antibody prepared from the primary antibody, the reporter antibody being conjugated to a detectable marker; and

detecting the reporter antibodies that bind to the captured digested microorganisms.

- 10 9. The method of claim 7 wherein the step of detecting comprises detecting the captured viability marker by detecting a change in a physical, a chemical, an optical, or an electrical property of the solid support.
 - 10. The method of claim 7 further comprising the steps of:

incubating the viability marker with a primary antibody specific for the
viability marker and conjugated to a reporter molecule, thereby forming a primary
antibody-antigen-reporter molecule sandwich; and

detecting the reporter molecule.

- 11. A method for detecting microorganisms according to the method of claim 1 comprising:
- 20 digesting the microorganisms;

incubating the digested microorganisms with a primary antibody specific for the viability marker;

conjugating the primary antibody to a reporter molecule to form a reporter-primary antibody complex; and

25 detecting the reporter molecule.

12. A method for detecting an actively respiring microorganisms in a sample comprising:

treating the microorganisms according to the method of claim 1; digesting the microorganisms;

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contacting a primary antibody prepared against a substituted formazan with the digested microorganisms;

contacting a secondary antibody prepared against the primary antibody, the secondary antibody being conjugated to a reporter molecule; and

detecting the reporter molecule.

- 13. The method of claim 12 further comprising the step of trapping the actively respiring microorganisms on a solid filtration membrane.
- 14. The method of claim 12 wherein the reporter molecule comprises an enzyme, a bioluminescent protein, a radioisotope, a chemiluminescent dye, a visible dye, a latex 10 particle, a magnetic particle or a fluorescent dye.
 - 15. The method of claim 12 wherein the sample is a clinical sample, a food sample, a cosmetic sample, a pharmaceutical sample, an industrial sample or an environmental sample.
 - 16. The method of claim 12 wherein the sample is a blood sample, a tissue sample, a tissue homogenate sample or a bodily fluid sample.
 - 17. The method of claim 12 wherein the microorganisms comprises a single species of microorganisms or a mixed population of microorganisms.
 - 18. The method of claim 12 wherein the sample contains less than 1000 cfu/mL.
 - 19. The method of claim 12 wherein the detecting takes less than two hours.
- 20 Monoclonal or polyclonal antibodies prepared to a substituted formazan and cross reactive to other formazans.
 - 21. A kit for the rapid and sensitive detection of viable microorganisms comprising: means for amplifying the presence of a microorganisms in a sample; and means for detecting the microorganisms.
- 25 22. A method for diagnosing a disease due to a microorganisms comprising: amplifying the presence of the microorganisms by the method of claim

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digesting the microorganisms;

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contacting a primary antibody prepared against a substituted formazan with the digested microorganisms;

contacting a secondary antibody prepared against the primary antibody, the secondary antibody being conjugated to a reporter molecule; and

detecting the reporter molecule.

23. A method for quantitating actively respiring microorganisms in a sample comprising:

contacting said microorganisms to a nutrient medium containing a predetermined amount of a tetrazolium salt;

10 metabolizing the tetrazolium salt to a viability marker using the microorganisms;

forming a quantitative amount of the viability marker that reflects the quantity of actively respiring microorganisms in the sample; and

detecting the viability marker.

24. A method for viability-marking an actively respiring microorganisms in a sample comprising contacting the contents of said sample to a nutrient medium containing a predetermined amount of a viability substrate, wherein metabolism of said viability substrate by the microorganisms of said sample produces a viability marker.